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From: Sent:	Russel, Jeffrey Friday, December	er 02, 2005 3:56 PM		
To: Subject:	STIC-Biotech/Cl Database Searc		•	
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Requester:				
_	Russel (TC1600)	)		
Art Unit: 1654				
Employee Num	ber:	$C \wedge$	. A	
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(9-50)				

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STRUCTURE FILE UPDATES: 11 DEC 2005 HIGHEST RN 869700-38-9 DICTIONARY FILE UPDATES: 11 DEC 2005 HIGHEST RN 869700-38-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

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\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* \* The CA roles and document type information have been removed from \* \* the IDE default display format and the ED field has been added, \* effective March 20, 2005. A new display format, IDERL, is now available and contains the CA role and document type information. \* +++++\*

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

7 S CCRECC/SQSP L1

ANSWER 1 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN L1

849178-34-3 REGISTRY RN

 $L-Cysteine, \ L-cysteinyl-L-cysteinyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-\alpha-glutamyl-L-arginyl-L-a$ CN cysteinyl- (9CI) (CA INDEX NAME)

43: PN: WO2005028615 SEQID: 58 unclaimed sequence SQL

1 CCRECC SEO =====

HITS AT: 1-6

\*\*RELATED SEQUENCES AVAILABLE WITH SEQLINK\*\*

REFERENCE 1: 142:368740

- ANSWER 2 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN L1
- RN 600706-61-4 REGISTRY
- L-Isoleucine, L-arginyl-L-valyl-L- $\alpha$ -aspartyl-L-alanyl-L-alanyl-L-CN alanyl-L-arginyl-L-\arginyl-L-alanyl-L-cysteinyl-L-cysteinyl-L-

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arginyl-L-\arglutamyl-L-cysteinyl-L-cysteinyl-L-alanyl-L-threonyl-
     L-alanyl- (9CI) (CA INDEX NAME)
OTHER NAMES:
    16: PN: WO03075856 SEQID: 16 unclaimed sequence
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SEQ
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           10-15
HITS AT:
REFERENCE
            1: 139:256227
    ANSWER 3 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN
L1
     439806-84-5 REGISTRY
RN
     L-Cysteine, L-cysteinyl-L-cysteinyl-L-arginyl-L-α-glutamyl-L-
CN
     cysteinyl-, cyclic 1,2:5,6-[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-
     1(3H),9'-[9H]xanthene]-4',5'-diyl)bis[arsonodithioite]] (9CI) (CA
     INDEX NAME)
SQL
    6
         1 CCRECC
SEQ
           =====
HITS AT:
           1-6
**RELATED SEQUENCES AVAILABLE WITH SEQLINK**
            1: 137:59787
REFERENCE
     ANSWER 4 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN
L1
     394709-23-0 REGISTRY
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     L-Isoleucine, L-arginyl-L-valyl-L-α-aspartyl-L-alanyl-L-alanyl-L-
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     arginyl-L-\alpha-glutamyl-L-cysteinyl-L-cysteinyl-L-alanyl-L-arginyl-
    L-alanyl- (9CI) (CA INDEX NAME)
OTHER NAMES:
    16: PN: W00210364 SEQID: 16 unclaimed sequence
CN
     59: PN: WO03075856 FIGURE: 4 unclaimed sequence
CN
SQL 19
         1 RVDAAAREAC CRECCARAI
SEO
                    = =====
           10-15
HITS AT:
REFERENCE
            1: 139:256227
REFERENCE
            2: 136:146127
    ANSWER 5 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN
L1
     268741-28-2 REGISTRY
RN
CN
     L-Alanine, L-tryptophyl-L-α-glutamyl-L-alanyl-L-alanyl-L-alanyl-
     L-arginyl-L-\alpha-glutamyl-L-alanyl-L-cysteinyl-L-cysteinyl-L-
     arginyl-L-\arglutamyl-L-cysteinyl-L-cysteinyl-L-alanyl-L-arginyl-
     (9CI) (CA INDEX NAME)
OTHER NAMES:
     46: PN: WO0114578 PAGE: 38 unclaimed sequence
CN
     4: PN: WO0047220 SEQID: 48 unclaimed sequence
CN
CN
     8: PN: US20040014071 SEQID: 4 unclaimed sequence
CN
     8: PN: WO2005038029 SEQID: 8 unclaimed sequence
CN
     9: PN: WO0153325 PAGE: 32 claimed sequence
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Shears

571-272-2528

Searcher

SQL 17

21

SEQ 1 WEAAAREACC RECCARA

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HITS AT: 9-14

\*\*RELATED SEQUENCES AVAILABLE WITH SEQLINK\*\*

REFERENCE 1: 142:425896

REFERENCE 2: 140:124793

REFERENCE 3: 135:149588

REFERENCE 4: 134:204756

REFERENCE 5: 133:172215

REFERENCE 6: 132:344976

L1 ANSWER 6 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN

RN 223673-79-8 REGISTRY

CN L-Alanine, L-alanyl-L-α-glutamyl-L-alanyl-L-alanyl-L-alanyl-Larginyl-L-α-glutamyl-L-alanyl-L-cysteinyl-L-cysteinyl-L-arginylL-α-glutamyl-L-cysteinyl-L-cysteinyl-L-alanyl-L-arginyl- (9CI)
(CA INDEX NAME)

OTHER NAMES:

CN 5: PN: WO0047220 SEQID: 49 unclaimed sequence

SQL 17

SEQ 1 AEAAAREACC RECCARA

== ====

HITS AT: 9-14

REFERENCE 1: 137:59787

REFERENCE 2: 133:172215

REFERENCE 3: 130:308804

L1 ANSWER 7 OF 7 REGISTRY COPYRIGHT 2005 ACS on STN

RN 223673-78-7 REGISTRY

CN L-Alaninamide, N-acetyl-L-tryptophyl-L-α-glutamyl-L-alanyl-L alanyl-L-alanyl-L-arginyl-L-α-glutamyl-L-alanyl-L-cysteinyl-L cysteinyl-L-arginyl-L-α-glutamyl-L-cysteinyl-L-cysteinyl-L alanyl-L-arginyl- (9CI) (CA INDEX NAME)

SQL 17

SEQ 1 WEAAAREACC RECCARA

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HITS AT: 9-14

\*\*RELATED SEQUENCES AVAILABLE WITH SEQLINK\*\*

REFERENCE 1: 137:59787

REFERENCE 2: 130:308804

FILE 'CAPLUS' ENTERED AT 12:36:39 ON 12 DEC 2005

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http://www.cas.org/infopolicy.html

L3 11 L1

L3 ANSWER 1 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 29 Apr 2005

ACCESSION NUMBER: 2005:371402 CAPLUS

DOCUMENT NUMBER: 142:425896

TITLE: Beetle luciferase reporter protein with various

modification motif increase or decrease

luminescence activity in the present or absent of

exogenous agent

INVENTOR(S): Fan, Frank; Lewis, Martin Ken; Schultz, John W.;

Wood, Keith V.; Butler, Braeden

PATENT ASSIGNEE(S): Promega Corporation, USA SOURCE: PCT Int. Appl., 178 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	PATENT NO.				KIND DATE			APPLICATION NO.						DATE		
					A2	-	2005	0428	1	WO 2	004-	 US32	705		2	0041001
							AU,									
			-	-			CZ,									
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		KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,
		MX,	MZ,	NA,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,
		SE,	SG,	SK,	SL,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	ŪG,	US,	UZ,
		VC,	VN,	YU,	ZA,	ZM,	ZW									
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	ŪG,	ZM,	ZW,
		AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,
		DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PL,
		PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,
		GW,	ML,	MR,	ΝE,	SN,	TD,	TG								
US	2005	1533	10		A1		2005	0714	,	US 2	004-	9574	33		2	0041001

PRIORITY APPLN. INFO.:

US 2003-510187P P 20031010

AB The current invention provides beetle luciferase reporter protein with various modifications. The reporter protein with modified motif in the absence or the present of an exogenous agent may enhance or inhibit luciferase activity.

IT 268741-28-2

RL: PRP (Properties)

(unclaimed sequence; beetle luciferase reporter protein with various modification motif increase or decrease luminescence activity in the present or absent of exogenous agent)

L3 ANSWER 2 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 01 Apr 2005

ACCESSION NUMBER: 2005:283572 CAPLUS

DOCUMENT NUMBER: 142:368740

TITLE: Plasmid vectors containing recombination sites and

topoisomerase recognition sites for detecting promoter activity and expressing fusion proteins Welch, Peter J.; Chesnut, Jonathan D.; Bennett,

Robert P.; Frimpong, Kenneth; Leong, Louis; Fan,

James; Yim, Harry; Vozza-Brown, Laura

PATENT ASSIGNEE(S): Invitrogen Corporation, USA

SOURCE: PCT Int. Appl., 378 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

INVENTOR(S):

	KIND	DATE	APF	LICAT	o. 	DATE			
			WO	2004-1	JS207	47		2	0040628
, CN, CO, , GD, GE, , KZ, LC,	CR, CU GH, GM LK, LR	, CZ, DE, , HR, HU, , LS, LT,	DK, DM ID, II LU, LV	I, DZ, I, IN, V, MA,	EC, I	EE, JP, MG,	EG, KE, MK,	ES, KG, MN,	FI, KP, MW,
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INFO.:			US	2003-	48730	1P		P 2	0030716
	615 615 , AG, AL, , CN, CO, , GD, GE, , KZ, LC, , MZ, NA, , SG, SK, , VN, YU, , GH, GM, , AZ, BY, , DK, EE, , RO, SE,	615 A2 615 A3 , AG, AL, AM, AT , CN, CO, CR, CU , GD, GE, GH, GM , KZ, LC, LK, LR , MZ, NA, NI, NO , SG, SK, SL, SY , VN, YU, ZA, ZM , GH, GM, KE, LS , AZ, BY, KG, KZ , DK, EE, ES, FI , RO, SE, SI, SK , ML, MR, NE, SN 615 A1	615 A2 20050331 615 A3 20050825 , AG, AL, AM, AT, AU, AZ, , CN, CO, CR, CU, CZ, DE, , GD, GE, GH, GM, HR, HU, , KZ, LC, LK, LR, LS, LT, , MZ, NA, NI, NO, NZ, OM, , SG, SK, SL, SY, TJ, TM, , VN, YU, ZA, ZM, ZW , GH, GM, KE, LS, MW, MZ, , AZ, BY, KG, KZ, MD, RU, , DK, EE, ES, FI, FR, GB, , RO, SE, SI, SK, TR, BF, , ML, MR, NE, SN, TD, TG 615 A1 20050505	615 A2 20050331 WO 615 A3 20050825  , AG, AL, AM, AT, AU, AZ, BA, BE , CN, CO, CR, CU, CZ, DE, DK, DM , GD, GE, GH, GM, HR, HU, ID, II , KZ, LC, LK, LR, LS, LT, LU, LV , MZ, NA, NI, NO, NZ, OM, PG, PH , SG, SK, SL, SY, TJ, TM, TN, TF , VN, YU, ZA, ZM, ZW , GH, GM, KE, LS, MW, MZ, NA, SI , AZ, BY, KG, KZ, MD, RU, TJ, TM , DK, EE, ES, FI, FR, GB, GR, HU , RO, SE, SI, SK, TR, BF, BJ, CE , ML, MR, NE, SN, TD, TG 615 A1 20050505 US INFO.:	615 A2 20050331 W0 2004-0615 A3 20050825  , AG, AL, AM, AT, AU, AZ, BA, BB, BG, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, GD, GE, GH, GM, HR, HU, ID, IL, IN, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, SG, SK, SL, SY, TJ, TM, TN, TR, TT, VN, YU, ZA, ZM, ZW  , GH, GM, KE, LS, MW, MZ, NA, SD, SL, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, DK, EE, ES, FI, FR, GB, GR, HU, IE, RO, SE, SI, SK, TR, BF, BJ, CF, CG, ML, MR, NE, SN, TD, TG  615 A1 20050505 US 2004-100505 US 2003-100505 US 2003-100505	615 A2 20050331 W0 2004-US207 615 A3 20050825  , AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, , CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, , GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, , KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, , MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, , SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, , VN, YU, ZA, ZM, ZW , GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, , AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, , DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, , RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, , ML, MR, NE, SN, TD, TG  615 A1 20050505 US 2003-48730  US 2003-48730	615 A2 20050331 W0 2004-US20747 615 A3 20050825  , AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, , CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, , GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, , KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, , MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, , SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, , VN, YU, ZA, ZM, ZW , GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, , AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, , DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, , RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, 615 A1 20050505 US 2003-487301P	615 A2 20050331 W0 2004-US20747 615 A3 20050825  , AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, , CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, , GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, , KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, , MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, , SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, , VN, YU, ZA, ZM, ZW , GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, , AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, , DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, , RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA,  ML, MR, NE, SN, TD, TG 615 A1 20050505 US 2003-487301P	615 A2 20050331 WO 2004-US20747 26 615 A3 20050825  , AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, , CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, , GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, , KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, , MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, , SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, , VN, YU, ZA, ZM, ZW , GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, , AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, , DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, , RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, , ML, MR, NE, SN, TD, TG 615 A1 20050505 US 2004-877952 2

AB The present invention provides nucleic acid mols. comprising one or more nucleic acid sequences encoding a polypeptide having a detectable activity, and in particular  $\beta$ -lactamase, said vectors comprising multiple recombination sites and/or topoisomerase recognition sites operably linked to a promoter. The present invention also provides methods of joining such nucleic acid mols. to nucleic acid mols. to be

assayed for promoter activity. The present invention also relates to methods of preparing fusion proteins comprising a polypeptide of interest and a polypeptide having a detectable activity. The GeneBLAzer System comprises the  $\beta$ -lactamase gene coupled with a fluorescence resonance energy transfer (FRET)-enabled substrate (CCF2, CCF2-FA, CCF2-AM, or CCF4-AM) and is an excellent reporter system for promoter studies in mammalian cells. A 'promoterless"  $\beta$ -lactamase vector (pGeneBlazer) may be constructed as a bidirectional TOPO vector, allowing PCR amplification of one or more promoters of interest and cloning of the promoters upstream of the  $\beta$ -lactamase gene. Recombination sites in combination with topoisomerase recognition sites allow joining of nucleic acids for expression of fusion proteins. Thus invention also uses nucleic acid regions encoding peptides with affinity for arsenic (Cys-Cys-X-X-Cys-Cys). The design, construction, and sequences of a variety of plasmid vectors is described.

#### IT 849178-34-3

RL: PRP (Properties)

(unclaimed sequence; plasmid vectors containing recombination sites and topoisomerase recognition sites for detecting promoter activity and expressing fusion proteins)

L3 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 23 Jan 2004

ACCESSION NUMBER: 2004:59670 CAPLUS

DOCUMENT NUMBER: 140:124793

TITLE: Methods for the detection, analysis and isolation of nascent proteins using non-radioactive markers

INVENTOR(S): Rothschild, Kenneth J.; Gite, Sadanand; Olejnik,

Jerzy; Lim, Mark

PATENT ASSIGNEE(S): Ambergen, Inc., USA

SOURCE: U.S. Pat. Appl. Publ., 147 pp., Cont.-in-part of

U.S. Ser. No. 49,332.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.			
	A1 20040122	US 2003-339712 US 1999-382736	20030110		
		WO 2000-US23233			
		BA, BB, BG, BR, BY, BZ,			
		DZ, EE, ES, FI, GB, GD,			
		JP, KE, KG, KP, KR, KZ,			
LR, LS, LT,	LU, LV, MA, MD,	MG, MK, MN, MW, MX, MZ,	NO, NZ,		
PL, PT, RO,	RU, SD, SE, SG,	SI, SK, SL, TJ, TM, TR,	TT, TZ,		
UA, UG, US,	UZ, VN, YU, ZA,	ZW, AM, AZ, BY, KG, KZ,	MD, RU,		
TJ, TM					
RW: GH, GM, KE,	LS, MW, MZ, SD,	SL, SZ, TZ, UG, ZW, AT,	BE, CH,		
CY, DE, DK,	ES, FI, FR, GB,	GR, IE, IT, LU, MC, NL,	PT, SE,		
BF, BJ, CF,	CG, CI, CM, GA,	GN, GW, ML, MR, NE, SN,	TD, TG		
US 2005009013	A1 20050113	US 2001-813197	20010320		
US 6875592	B2 20050405				
US 2003190643	A1 20031009	US 2002-264127	20021003		
	A1 20050210	US 2003-719523	20031121		
CA 2512552	AA 20040729	CA 2004-2512552	20040109		

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20040729
                                           WO 2004-US528
                                                                  20040109
    WO 2004063714
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            BB, BG, BG, BR, BR, BW, BY, BY, BZ, BZ, CA, CH, CN, CN, CO,
            CO, CR, CR, CU, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EC, EC,
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            MA, MD, MD, MG, MK, MN, MW, MX, MX, MZ
                              20051005
                                          EP 2004-701238
                         A2
    EP 1581797
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
            PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
PRIORITY APPLN. INFO.:
                                           US 1999-382736
                                                               A2 19990825
                                           WO 2000-US23233
                                                               W 20000823
                                                               A2 20020621
                                           US 2002-49332
                                           US 1999-382950
                                                               A 19990825
                                           US 2001-813197
                                                               A1 20010320
                                           US 2003-339712
                                                               A 20030110
                                           WO 2004-US528
                                                               W 20040109
```

This invention relates to non-radioactive markers that facilitate the detection and anal. of nascent proteins translated within cellular or cell-free translation systems. Nascent proteins containing these markers can be rapidly and efficiently detected, isolated and analyzed without the handling and disposal problems associated with radioactive reagents. Preferred markers are dipyrrometheneboron difluoride (4,4-difluoro-4-bora-3a,4a-diaza-s-indacene) dyes.

## IT 268741-28-2

RL: PRP (Properties)

(unclaimed sequence; methods for the detection, anal. and isolation of nascent proteins using non-radioactive markers)

L3 ANSWER 4 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 19 Sep 2003

ACCESSION NUMBER: 2003:737532 CAPLUS

DOCUMENT NUMBER: 139:256227

DOCUMENT NUMBER: 139.230227

TITLE: Methods for enhancing oligonucleotide-directed nucleic acid sequence alteration using repair proteins, histone deacetylase inhibitors,  $\lambda$  phage  $\beta$  proteins and hydroxyurea for use in

therapy of blood diseases

INVENTOR(S): Kmiec, Eric B.; Parekh-Olmedo, Hetal; Brachman,

Erin E.

PATENT ASSIGNEE(S): University of Delaware, USA

SOURCE: PCT Int. Appl., 135 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003075856	A2	20030918	WO 2003-US7217	20030307

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WO 2003075856
                          A3
                                20040624
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ,
             LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ,
             NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM,
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE,
             SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
             NE, SN, TD, TG
                                            CA 2003-2478479
                                20030918
                                                                    20030307
    CA 2478479
                          AA
    US 2003207451
                                            US 2003-384918
                                                                    20030307
                          A1
                                20031106
                                            EP 2003-716412
                                                                    20030307
                          A2
                                20041229
    EP 1490013
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
             PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                           JP 2003-574132
                                                                    20030307
                         T2
                                20050630
     JP 2005518817
                                            US 2002-363053P
                                                                P 20020307
PRIORITY APPLN. INFO.:
                                                                   20020307
                                            US 2002-363054P
                                                                Р
                                            US 2002-363341P
                                                                Ρ
                                                                    20020307
                                            US 2002-416983P
                                                                    20021007
                                            WO 2003-US7217
                                                                W
                                                                   20030307
     Improved methods, compns., and kits for oligonucleotide-mediated
AB
     nucleic acid sequence alteration using repair proteins, histone
     deacetylase inhibitors and hydroxyurea are provided. These methods
     may be use for treatment of blood disorders.
     394709-23-0 600706-61-4
IT
     RL: PRP (Properties)
        (unclaimed sequence; methods for enhancing oligonucleotide-directed
        nucleic acid sequence alteration using repair proteins, histone
        deacetylase inhibitors, \lambda phage \beta proteins and
        hydroxyurea for use in therapy of blood diseases)
    ANSWER 5 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN
    Entered STN: 02 May 2002
                         2002:326835 CAPLUS
ACCESSION NUMBER:
                         137:59787
DOCUMENT NUMBER:
                         New biarsenical ligands and tetracysteine motifs
TITLE:
                         for protein labeling in vitro and in vivo:
                         Synthesis and biological applications
                         Adams, Stephen R.; Campbell, Robert E.; Gross,
AUTHOR(S):
                         Larry A.; Martin, Brent R.; Walkup, Grant K.; Yao,
                         Yong; Llopis, Juan; Tsien, Roger Y.
                         Department of Pharmacology, Department of
CORPORATE SOURCE:
                         Chemistry and Biochemistry, Howard Hughes Medical
                         Institute and Biomedical Sciences Program,
                         University of California San Diego, La Jolla, CA,
                         92093-0647, USA
                         Journal of the American Chemical Society (2002),
SOURCE:
                         124(21), 6063-6076
                         CODEN: JACSAT; ISSN: 0002-7863
                         American Chemical Society
PUBLISHER:
DOCUMENT TYPE:
                         Journal
```

LANGUAGE: English

We recently introduced a method (Griffin, B. A.; Adams, S. R.; Tsien, R. Y. Science 1998, 281, 269-272 and Griffin, B. A.; Adams, S. R.; Jones, J.; Tsien, R. Y. Methods Enzymol. 2000, 327, 565-578) for site-specific fluorescent labeling of recombinant proteins in living cells. The sequence Cys-Cys-Xaa-Xaa-Cys-Cys, where Xaa is an noncysteine amino acid, is genetically fused to or inserted within the protein, where it can be specifically recognized by a membrane-permeant fluorescein derivative with two As(III) substituents, Flash, which fluoresces only after the arsenics bind to the cysteine thiols. We now report kinetics and dissociation consts. (.apprx.10-11 M) for FlAsH binding to model tetracysteine peptides. Affinities in vitro and detection limits in living cells are optimized with Xaa-Xaa = Pro-Gly, suggesting that the preferred peptide conformation is a hairpin rather than the previously proposed  $\alpha$ -helix. Many analogs of FlAsH have been synthesized, including ReAsH, a resorufin derivative excitable at 590 nm and fluorescing in the red. Analogous biarsenicals enable affinity chromatog., fluorescence anisotropy measurements, and electron-microscopic localization of tetracysteine-tagged proteins.

## IT 223673-78-7 223673-79-8 439806-84-5

RL: ARU (Analytical role, unclassified); ANST (Analytical study) (biarsenical ligands and tetracysteine motifs for protein labeling in vitro and in vivo)

REFERENCE COUNT:

THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 6 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

45

Entered STN: 10 Feb 2002

ACCESSION NUMBER: 2002:107524 CAPLUS

DOCUMENT NUMBER: 136:146127

TITLE: Methods for enhancing targeted gene alteration in

cells having altered activity of DNA repair proteins using chimeric RNA-DNA double-stranded

hairpin oligonucleotides

INVENTOR(S): Kmiec, Eric B.; Gamper, Howard B.; Rice, Michael

C.; Liu, Li

PATENT ASSIGNEE(S): University of Delaware, USA

SOURCE: PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

PATENT NO.					KIND DATE			APPLICATION NO.						DATE		
	2002				A2 A3		2002 2003		1	WO 2	001-	US23	770		2	0010727
".0			AG,						BA.	BB.	BG,	BR,	BY,	BZ,	CA,	CH,
		•	co,	•	•		•	•	-	-	-	-				
			GH,	•		•	-	-	-	-		-	-	-	-	
		-	LK,													
		NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,
		TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	zw					
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪG,	ZW,	AM,	ΑZ,	BY,
		KG,	ΚZ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,
		GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	ВJ,	CF,	CG,	CI,

CA 24173 EP 13640 R:	08 AT, BE, C	AA A2 H, DE,	2002 2003 DK, ES,	0207 1126 FR,	CA EP GB, GI	2001- 2001- R, IT,	2417344 957311 LI, LU,		20010727 20010727 E, MC,
	PT, IE, S								20020720
	17377								20020730
US 20032	15947	Al	2003	1120					20030124
PRIORITY APPL	N. INFO.:				US	2000-	220999P	P	20000727
					US	2000-	244989P	P	20001030
					US	2000-	192176P	P	20000327
					US	2000-	192179P	P	20000327
					US	2000-	208538P	P	20000601
					US	2001-	818875	A3	20010327
					WO	2001-	us23770	W	20010727

AB Methods are presented for enhancing the efficiency of oligonucleotide-mediated repair or alteration of genetic information in cells having altered activity of DNA repair proteins using chimeric RNA-DNA double-stranded. The methods comprise using cells or cell-free exts. having altered levels or activity of at least one protein from the RAD52 epistasis group, the mismatch repair group or the nucleotide excision repair group. A assay system for identifying inhibitors of DNA repair proteins and monitoring genetic alteration using the oligonucleotides of the invention is also presented. Kits comprising cells and cell-free exts. having reduced activity of DNA repair proteins and vectors for enhancing targeted gene alteration are also presented. The invention demonstrates that gene repair depends on the dose of DNA repair proteins and expression of RAD52 gene suppresses oligonucleotide-directed gene alteration.

## IT 394709-23-0

RL: PRP (Properties)

(unclaimed sequence; methods for enhancing targeted gene alteration in cells having altered activity of DNA repair proteins using chimeric RNA-DNA double-stranded hairpin oligonucleotides)

L3 ANSWER 7 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 27 Jul 2001

ACCESSION NUMBER: 2001:545718 CAPLUS

DOCUMENT NUMBER: 135:149588

TITLE: Method of affinity purifying proteins using

modified bis-arsenical fluorescein

INVENTOR(S): Vale, Ronald D.; Thorn, Kurt; Cooke, Roger;

Matuska, Marija; Naber, Nariman

Matuska, Marija; Naper, Nariman

PATENT ASSIGNEE(S): The Regents of the University of California, USA

PATENT ASSIGNEE(S): The Regents of the Uni SOURCE: PCT Int. Appl., 52 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE

WO	2001	05332	25		A2			0726	W	0	200	1-U	JS22	14		2	0010	122
WO	2001	05332	25		A3	:	2002	0307										
	W:	AU,	CA,	JP														
	RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR	, G	В,	GR,	ΙE,	IT,	LU,	MC,	
		NL,	PT,	SE,	TR													
US	6831	160			В1	:	2004	1214	U	S	200	0-5	0266	54		2	00002	211
AU	2001	0310	36		A5	:	2001	0731	А	U	200	1-3	31086	5		2	0010	L22
US	2005	23342	28		<b>A1</b>	:	2005	1020	U	S	200	4-1	.2853	3		2	00412	214
PRIORITY	APP	LN.	INFO	.:					ប	S	200	0-1	.7805	54P	;	P 2	0000	124
									U	S	200	0-5	0266	54	7	A. 2	00002	211
									W	0	200	1-U	JS221	l 4	1	N 2	0010	L22

OTHER SOURCE(S): MARPAT 135:149588

AB The present invention features methods for purifying polypeptides of interest using a modified Fluorescein arsenical helix binder (FlAsH) compound immobilized on a solid support. An exemplary FlAsH target sequence motif is also presented. Examples of modification of the FlAsH compound which allow immobilization to a solid support are also provided. The present invention also provides DNA constructs for producing a dual affinity tagged polypeptide and methods for purification thereof. Human kinesin constructs C-terminally tagged with the peptide WEAAAREACCRECCARA (specifically chelating with  $\beta$ -alanine-modified FlAsH, preparation given) were expressed in Escherichia coli and purified using beads containing  $\beta$ -alanine-modified FlAsH. Protein was eluted using 1,2-ethanedithiol.

IT 268741-28-2P

RL: BPN (Biosynthetic preparation); BPR (Biological process); BSU (Biological study, unclassified); NUU (Other use, unclassified); PRP (Properties); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(amino acid sequence, as FlAsH peptide target; affinity purifying proteins using modified bis-arsenical fluorescein)

L3 ANSWER 8 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 02 Mar 2001

ACCESSION NUMBER: 2001:152863 CAPLUS

DOCUMENT NUMBER: 134:204756

TITLE: Methods for the detection, analysis and isolation

of nascent proteins

INVENTOR(S): Rothschild, Kenneth J.; Gite, Sadanand; Olejnik,

Jerzy

PATENT ASSIGNEE(S): Ambergen, Inc., USA SOURCE: PCT Int. Appl., 204 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.				KIND DATE			;	APPL:		DATE					
WO 20010				A1 20010301								_			
W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,
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	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,
	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,
	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,

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         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                           US 1999-382950
                                20011016
    US 6303337
                          B1
                                20011023
                                            US 1999-382736
    US 6306628
                          B1
                                20010301
                                            CA 2000-2383554
                                                                   20000823
    CA 2383554
                         AA
                                            EP 2000-957758
                                                                   20000823
                                20020605
    EP 1210449
                         A1
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
            PT, IE, SI, LT, LV, FI, RO, MK, CY, AL
                                            JP 2001-518889
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                                20040513
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    AU 775940
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                                20040819
                                            AU 2000-69328
                                20050113
                                            US 2001-813197
                                                                   20010320
    US 2005009013
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    US 6875592
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                                20050405
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    US 2002132248
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    US 2003092031
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    US 2003190643
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                         A1
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    US 2004014071
                                                                   20031121
                               20050210
                                            US 2003-719523
                         A1
    US 2005032078
                                                                A 19990825
                                            US 1999-382736
PRIORITY APPLN. INFO.:
                                            US 1999-382950
                                                                A 19990825
                                            WO 2000-US23233
                                                                W 20000823
                                            US 2001-813197
                                                                A1 20010320
                                                                A2 20020621
                                            US 2002-49332
    This invention relates to non-radioactive markers that facilitate the
AΒ
    detection and anal. of nascent proteins translated within cellular or
    cell-free translation systems. Nascent proteins containing these markers
    can be rapidly and efficiently detected, isolated and analyzed without
    the handling and disposal problems associated with radioactive reagents.
    Preferred markers are dipyrrometheneboron difluoride
     (4,4-difluoro-4-bora-3a,4a-diaza-s-indacene) dyes.
IT
    268741-28-2
    RL: PRP (Properties)
        (unclaimed sequence; methods for the detection, anal. and isolation
        of nascent proteins)
                               THERE ARE 4 CITED REFERENCES AVAILABLE FOR
REFERENCE COUNT:
                         4
                               THIS RECORD. ALL CITATIONS AVAILABLE IN THE
                               RE FORMAT
    ANSWER 9 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN
L3
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ED Entered STN: 18 Aug 2000

ACCESSION NUMBER: 2000:573678 CAPLUS

DOCUMENT NUMBER: 133:172215

TITLE: Controlling protein levels in eucaryotic organisms

using novel compds. comprising a ubiquitination recognition element and a protein binding element

INVENTOR(S): Kenten, John H.; Roberts, Steven F.; Lebowitz,

Michael S.

PATENT ASSIGNEE(S): Proteinix, Inc., USA SOURCE: PCT Int. Appl., 106 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PA'						rińD		DATE		APPLICATION NO.					DATE	
WO.	2000	0472	20		A1	_	2000	0817		wo	2000-	US34:	36			20000211
	W:										, BR,					
											, GH,					
		IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC	, LK,	LR,	LS,	LT,	LU	, LV,
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	$\mathtt{PL}$	, PT,	RO,	RU,	SD,	SE	, SG,
		SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	UA,	ŬĠ	, US,	UΖ,	VN,	YU,	ZA	, ZW,
							MD,									
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ	, UG,	ZW,	ΑT,	BE,	CH	, CY,
																, BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML	, MR,	ΝE,	SN,	TD,	TG	
US	6306	663			B1		2001	1023		US	1999-	4067	81			19990928
CA	2362	560			AA		2000	0817		CA	2000-	2362	560			20000211
EP	1156	817			A1		2001	1128		ΕP	2000-	9085	80			20000211
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE	, MC,
		PT,	ΙE,	SI,	LT,	LV,	FI,	RO								
JP	2002	5364	17		Т2		2002	1029			2000-					20000211
US	2002	1468	43		A1		2002	1010								20010614
US	2002	1730	49		A1		2002	1121		US	2001-	8801	32			20010614
	6559						2003									
US	2003	1537	27		<b>A</b> 1		2003	0814					-			20030116
IORIT	Y APP	LN.	INFO	.:						US	1999-	1198	51P		P	19990212
										US	1999-	4067	81		A2	19990928
										WO	2000-	US34	36		W	20000211
										US	2001-	8801	32	(,	<b>A</b> 3	20010614

The invention relates to novel compds. comprising a ubiquitination recognition element and a protein binding element. The invention also relates to the use of said compds. for modulating the level and/or activity of a target protein. The compds. are useful for the treatment of diseases such as infections, inflammatory conditions, cancer and genetic diseases. The compds. are also useful as insecticides and herbicides.

IT 223673-79-8 268741-28-2

RL: PRP (Properties)

(unclaimed sequence; controlling protein levels in eucaryotic organisms using novel compds. comprising a ubiquitination recognition element and a protein binding element)

REFERENCE COUNT:

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 10 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

9

ED Entered STN: 16 Mar 2000

ACCESSION NUMBER: 2000:169797 CAPLUS

DOCUMENT NUMBER: 132:344976

TITLE: A novel method of affinity-purifying proteins

using a bis-arsenical fluorescein

AUTHOR(S): Thorn, Kurt S.; Naber, Nariman; Matuska, Marija;

Vale, Ronald D.; Cooke, Roger

CORPORATE SOURCE: Department of Cellular and Molecular Pharmacology,

University of California, San Francisco, CA,

94143, USA

SOURCE: Protein Science (2000), 9(2), 213-217

CODEN: PRCIEI; ISSN: 0961-8368

PUBLISHER: Cambridge University Press

DOCUMENT TYPE: Journal LANGUAGE: English

Genetically-encoded affinity tags constitute an important strategy for purifying proteins. Here, we have designed a novel affinity matrix based on the bis-arsenical fluorescein dye FlAsH, which specifically recognizes short  $\alpha$ -helical peptides containing the sequence CCXXCC. We find that kinesin tagged with this cysteine-containing helix binds specifically to FlAsH resin and can be eluted in a fully active form. This affinity tag has several advantages over polyhistidine, the only small affinity tag in common use. The protein obtained with this single chromatog. step from crude Escherichia coli lysates is purer than that obtained with nickel affinity chromatog. of 6xHis tagged kinesin. Moreover, unlike nickel affinity chromatog., which requires high concns. of imidazole or pH changes for elution, protein bound to the FlAsH column can be completely eluted by dithiothreitol. Because of these mild elution conditions, FlAsH affinity chromatog. is ideal for recovering fully active protein and for the purification of intact protein complexes.

IT 268741-28-2

RL: BPR (Biological process); BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); PROC (Process); USES (Uses)

(novel method of affinity-purifying proteins using a bis-arsenical fluorescein)

REFERENCE COUNT:

THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 11 OF 11 CAPLUS COPYRIGHT 2005 ACS on STN

ED Entered STN: 10 May 1999

ACCESSION NUMBER: 1999:286159 CAPLUS

DOCUMENT NUMBER:

130:308804

TITLE:

Target protein sequences for binding of synthetic

biarsenical molecules

INVENTOR(S):

Tsien, Roger Y.; Griffin, Albert B.

PATENT ASSIGNEE(S):

The Regents of the University of California, USA

SOURCE: PCT Int. Appl., 77 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	PATENT NO.				KIND DATE				APPLICATION NO.						DATE		
WO	9921	013			A1 19990429			WO 1	998-	US22:	363		1:	9981021			
	W:	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	
		DE,	DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IS,	JP,	
		KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	
		MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	
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	RW:	GH,															
		ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	
		CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG				

US US	59324 60083 60542 99113	378 271			A A A A1	1999 2000	0803 1228 0425 0510	us us	1997- 1997-	955206 955859 955050 11139		1 1	.9971021 .9971021 .9971021
	1032				A1		0906			953881			9981021
	R:	AT,	BE, IE,		-	DK, ES,		GB, GI	R, IT,	LI, LU	, NL,	SE,	MC,
US	6451	•			В1	2002	0917	US	1999-	372338		1	.9990811
US	2003	0833	73		A1	2003	0501	US	2002-	126752		2	0020419
US	6686	458			B2	2004	0203						
US	2005	1312	17		A1	2005	0616	US	2004-	772164		2	0040203
PRIORITY	APP	LN. I	INFO	.:				US	1997-	955050	A	2 1	.9971021
								US	1997-	955206	A	.2 1	.9971021
								US	1997-	955859	P	2 1	9971021
								WO	1998-	US22363	W	1	.9981021
								US	1999-	372338	P	.1 1	.9990811
								US	2002-	126752	A	.1 2	0020419

OTHER SOURCE(S): MARPAT 130:308804

The present invention features biarsenical mols. and target sequences that specifically react with the biarsenical mols. A bonding partner comprises a carrier polypeptide and a target sequence, wherein the target sequence is heterologous to the carrier polypeptide and the target sequence contains one or more cysteines capable of specifically reacting with a biarsenical mol. Bonding partners that include target sequences, vectors that include nucleic acid sequences that encode the target sequences and host cells that include the target sequences are also featured in the invention. One example of a biarsenical compound is an arsenical derivative of fluorescein.

# IT 223673-78-7

RL: ARU (Analytical role, unclassified); BPR (Biological process); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); PROC (Process)

(SEQ ID 1; target protein sequences for binding of synthetic biarsenical mols.)

#### IT 223673-79-8

RL: ARU (Analytical role, unclassified); BPR (Biological process); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); PROC (Process)

(SEQ ID 4; target protein sequences for binding of synthetic biarsenical mols.)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

FILE 'MEDLINE' ENTERED AT 12:36:52 ON 12 DEC 2005

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L4 0 L1

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OM protein - protein search, using sw model

Run on: December 8, 2005, 15:48:28; Search time 229 Seconds

(without alignments)

52.375 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2166443 seqs, 705528306 residues

Total number of hits satisfying chosen parameters: 2166443

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : UniProt\_05.80:\*

1: uniprot\_sprot:\*
2: uniprot trembl:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

		용					
Result		Query					
No.	Score	Match	Length	DB	ID	Descrip	ption
1	53	52.5	157	1	VE6_HPV12	P36803	human papil
2	51	50.5	278	2	Q61GH0_DROME	Q6igh0	drosophila
3	51	50.5	502	2	Q9BGM9_9MAMM	Q9bgm9	tachyglossu
4	50	49.5	1370	1	ZN261 HUMAN	Q14202	homo sapien
5	50	49.5	1370	1	ZN261 MOUSE	Q9jlm4	mus musculu
6	49.5	49.0	602	2	Q75NZ5_CHLRE	Q75nz5	chlamydomon
7	49	48.5	115	1	ALK1 PIG	P22298	sus scrofa
8	49	48.5	155	2	Q9PXB1_HPV08	Q9pxb1	human papil
9	49	48.5	168	1	VE6_HPV21	P28832	human papil
10	49	48.5	1067	2	Q4QFE4_LEIMA		leishmania
11	48.5	48.0	390	2	Q4S604_TETNG	Q4s604	tetraodon n
12	48	47.5	62	2	Q4PN38_IXOSC	Q4pn38	ixodes scap
13	48	47.5	131	1	ALK1_MOUSE	P97430	mus musculu
14	48	47.5	131	2	Q548X8_MOUSE	Q548x8	mus musculu
15	48	47.5	157	2	040617_HPVR7	040617	human papil

16	48	47.5	157	2	Q81986_HPV05	Q81986	human papil
17	48	47.5	157	2	Q913V6 9PAPI	Q913v6	human papil
18	47.5	47.0	525	2	Q64FQ2 ARATH	Q64fq2	arabidopsis
19	47.5	47.0	676	2	048785 ARATH	048785	arabidopsis
20	47	46.5	88	2	Q62H93_BURMA	Q62h93	burkholderi
21	47	46.5	101	2	Q4IVT4 AZOVI		azotobacter
22	47	46.5	131	2	Q9R0Z8 RAT		rattus norv
23	47	46.5	156	1	VE6 HPV47		human papil
24	47	46.5	171	1	VE6 HPV14		human papil
25	47	46.5	181	2	Q8VMH1 PSEPU		pseudomonas
26	47	46.5	193	1	KR415 HUMAN		homo sapien
	47	46.5	210	1	KRA47 HUMAN		homo sapien
27					_		pseudomonas
28	47	46.5	219	2	Q52396_PSEST		
29	47	46.5	1175	2	Q4P5X7_USTMA		ustilago ma
30	46	45.5	80	1	IBB4_LONCA		lonchocarpu
31	46	45.5	88	2	Q52509_PSESX		pseudomonas
32	46	45.5	129	1	KRA56_HUMAN		homo sapien
33	46	45.5	161	2	Q8MZ55_DROME		drosophila
34	46	45.5	166	1	VE6_HPV19		human papil
35	46	45.5	186	1	KRA45_HUMAN	=	homo sapien
36	46	45.5	191	2	Q28583_SHEEP	Q28583	ovis aries
37	46	45.5	298	2	Q65T35_MANSM	Q65t35	mannheimia
38	46	45.5	412	2	P91666_DROME	P91666	drosophila
39	46	45.5	465	1	HYIN2 BRAJA	P19922	bradyrhizob
40	46	45.5	491	2	Q4T2B4 TETNG	Q4t2b4	tetraodon n
41	46	45.5	757	2	Q6PFS4 BRARE		brachydanio
42	46	45.5	1033	2	Q4T6W6 TETNG		tetraodon n
43	46	45.5	1063	2	Q4TBG6 TETNG		tetraodon n
44	46	45:5	1367	2	Q629H4_CAEBR	_	caenorhabdi
45	46	45.5	1376	2	Q23590 CAEEL		caenorhabdi
46	46	45.5	1955	2	Q9VXG2_DROME		drosophila
47	46	45.5	1959	2	Q9VXG1 DROME		drosophila
48	45.5	45.0	139	2	Q8RYZ5_ORYSA		oryza sativ
49	45.5	45.0	279	2	Q4RZU3_TETNG		tetraodon n
			61	2			
50	45	44.6		2	Q9PB81_XYLFA		xylella fas
51	45	44.6	68		097751_PIG		sus scrofa
52	45	44.6	100	2	Q5UPC9_MIMIV	· -	mimivirus.
53	45	44.6	117	2	Q76YA2_9CAUD	· · · · · · · · · · · · · · · · · · ·	bacteriopha
54	45	44.6	120	2	Q9QQ85_HPV08		human papil
55	45	44.6	130	2	Q6IE20_RAT		rattus norv
56	45	44.6	149	2	O12671_9PAPI		colobus mon
57	45	44.6	155	1	VE6_HPV08		human papil
58	45	44.6	157	1	VE6_HPV05		human papil
59	45	44.6	157	1	VE6_HPV36		human papil
60	45	44.6	157	1	VE6_HPV5B		human papil
61	45	44.6	157	2	Q76WJ7_HPV5B	Q76wj7	human papil
62	45	44.6	157	2	Q6LBH6_HPV05	Q61bh6	human papil
63	45	44.6	157	2	Q6YNY6_9PAPI	Q6yny6	human papil
64	45	44.6	157	2	Q81962 HPV05	Q81962	human papil
65	45	44.6	157	2	Q81985 HPV05	Q81985	human papil
66	45	44.6	157	2	Q910D7 9PAPI		human papil
67	45	44.6	157	2	Q910X3 9PAPI		human papil
68	45	44.6	157	2	Q913V7 9PAPI		human papil
69	45	44.6	157	2	Q913V8 9PAPI		human papil
70	45	44.6	157		Q913V9 9PAPI		human papil
71	45	44.6	157	2	Q913W0 9PAPI		human papil
72	45	44.6	157	2	Q913W1 9PAPI		human papil
				_		Q213W1	papir

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45 44.6 157 2 Q913W2 9PAPI
                                                              Q913w2 human papil
    73
            45 44.6 157 2 Q913W3 9PAPI
                                                             Q913w3 human papil
    74
            45 44.6 157 2 Q913W4 9PAPI
                                                             Q913w4 human papil
    75
            45 44.6 157 2 Q913W4_9PAPI

45 44.6 157 2 Q913W5_9PAPI

45 44.6 157 2 Q913W6_9PAPI

45 44.6 165 1 VE6_HPV20

45 44.6 165 2 Q9D7P3_MOUSE

45 44.6 233 2 Q7RZM5_NEUCR
                                                             Q913w5 human papil
Q913w6 human papil
    76
    77
P28831 human papil
    78
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RESULT 1
VE6 HPV12
                STANDARD;
                              PRT; 157 AA.
   VE6 HPV12
AC
    P36803;
    01-JUN-1994 (Rel. 29, Created)
DT
DT 01-JUN-1994 (Rel. 29, Last sequence update)
DT 13-SEP-2005 (Rel. 48, Last annotation update)
DE E6 protein.
GN
    Name=E6;
os
    Human papillomavirus type 12.
OC
    Viruses; dsDNA viruses, no RNA stage; Papillomaviridae;
OC
    Betapapillomavirus.
OX
    NCBI TaxID=10604;
RN
    [1]
    NUCLEOTIDE SEQUENCE [GENOMIC DNA].
RP
    MEDLINE=94265501; PubMed=8205838;
RX
RA
    Delius H., Hofmann B.;
    "Primer-directed sequencing of human papillomavirus types.";
RT
   Curr. Top. Microbiol. Immunol. 186:13-31(1994).
RL
CC
    -!- FUNCTION: Transcriptional transactivator. Binds double stranded
CC
       DNA (By similarity).
CC
   -!- SUBCELLULAR LOCATION: Nuclear matrix-associated (By similarity).
CC
    -!- SIMILARITY: Belongs to the papillomaviruses E6 protein family.
    CC
```

```
This Swiss-Prot entry is copyright. It is produced through a collaboration
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
    the European Bioinformatics Institute. There are no restrictions on its
CC
    use as long as its content is in no way modified and this statement is not
CC
CC
    removed.
    _____
CC
    EMBL; X74466; CAA52496.1; -; Genomic DNA.
DR
    PIR; S36538; S36538.
    InterPro; IPR001334; E6.
DR
DR
    Pfam; PF00518; E6; 1.
KW
    Activator; DNA-binding; Early protein; Metal-binding; Nuclear protein;
KW
    Transcription; Transcription regulation; Zinc; Zinc-finger.
                             Potential.
              39 . 75
FT
    ZN FING
FT
    ZN FING
               112
                     148
                              Potential.
SO
    SEOUENCE
              157 AA; 17984 MW; E9EC735537733FDC CRC64;
                        52.5%; Score 53; DB 1; Length 157;
 Query Match
 Best Local Similarity 53.3%; Pred. No. 12;
 Matches
           8; Conservative
                              1; Mismatches
                                                6; Indels 0; Gaps
                                                                         0;
          1 WEAAAREACCRECCA 15
Qу
                   |:
Db
          63 WKGHFVTACCRSCCA 77
RESULT 2
O6IGHO DROME
ID
    Q6IGHO DROME PRELIMINARY; PRT;
                                        278 AA.
AC
    05-JUL-2004 (TrEMBLrel. 27, Created)
DT
    05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
DT
    05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
DT
DE
    HDC06306.
GN
    ORFNames=HDC06306;
OS
    Drosophila melanogaster (Fruit fly).
    Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
    Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
OC
OC
    Ephydroidea; Drosophilidae; Drosophila.
OX
    NCBI TaxID=7227;
RN
    NUCLEOTIDE SEQUENCE.
RP
    PubMed=14709175; DOI=10.1186/gb-2003-5-1-r3;
RX
    Hild M., Beckmann B., Haas S.A., Koch B., Solovyev V., Busold C.,
RA
RA
    Fellenberg K., Boutros M., Vingron M., Sauer F., Hoheisel J.D.,
    Paro R.;
RA
RT
    "An integrated gene annotation and transcriptional profiling approach
RT
    towards the full gene content of the Drosophila genome.";
    Genome Biol. 5: RESEARCH0003.1-RESEARCH0003.17(2003).
RL
CC
    -!- MISCELLANEOUS: The sequence shown here is derived from an
CC
        EMBL/GenBank/DDBJ third party annotation (TPA) entry.
DR
    EMBL; BK003796; DAA02494.1; -; Genomic DNA.
DR
    InterPro; IPR006209; EGF like.
DR
    PROSITE; PS00022; EGF 1; UNKNOWN 1.
SO
    SEQUENCE 278 AA; 32016 MW; 06E7253102FE5BF1 CRC64;
 Query Match
                        50.5%; Score 51; DB 2; Length 278;
 Best Local Similarity 87.5%; Pred. No. 35;
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CC

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7; Conservative 0; Mismatches 1; Indels
                                                                0; Gaps
                                                                            0;
 Matches
           9 CCRECCAR 16
Qу
              250 CCRECCCR 257
Db
RESULT 8
Q9PXB1 HPV08
     O9PXB1 HPV08 PRELIMINARY;
                                     PRT;
                                           155 AA.
ID
AC
     O9PXB1;
     01-MAY-2000 (TrEMBLrel. 13, Created)
DT
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT
     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DT
DΕ
     E6 protein.
     Human papillomavirus type 8.
OS
     Viruses; dsDNA viruses, no RNA stage; Papillomaviridae;
OC
OC
     Papillomavirus.
     NCBI TaxID=10579;
OX
RN
     [1]
     NUCLEOTIDE SEQUENCE.
RP
RX
     MEDLINE=91361540; PubMed=1653484;
     Deau M.C., Favre M., Orth G.;
RA
     "Genetic heterogeneity among human papillomaviruses (HPV) associated
RT
     with epidermodysplasia verruciformis: evidence for multiple allelic
RT
     forms of HPV5 and HPV8 E6 genes.";
RT
     Virology 184:492-503(1991).
RL
DR
     GO; GO:0042025; C:host cell nucleus; IEA.
     GO; GO:0005634; C:nucleus; IEA.
DR
     GO; GO:0003677; F:DNA binding; IEA.
DR
     GO; GO:0006355; P:regulation of transcription, DNA-dependent; IEA.
DR
     InterPro; IPR001334; E6.
DR
     Pfam; PF00518; E6; 1.
DR
     SEOUENCE
              155 AA; 17764 MW; 6986A0F88C7A33FD CRC64;
SO
                          48.5%; Score 49; DB 2; Length 155;
  Query Match
                          53.3%; Pred. No. 41;
  Best Local Similarity
                                1; Mismatches
                                                6; Indels
            8; Conservative
                                                                0; Gaps
  Matches
                                                                             0;
            1 WEAAAREACCRECCA 15
QУ
              :
                     63 WKNYVVTACCRCCCA 77
Db
RESULT 30
IBB4 LONCA
     IBB4 LONCA
                    STANDARD;
                                   PRT;
                                           80 AA.
ID
AC
     P16343;
     01-AUG-1990 (Rel. 15, Created)
DT
     01-AUG-1990 (Rel. 15, Last sequence update)
DT
     10-MAY-2005 (Rel. 47, Last annotation update)
DT
     Bowman-Birk type proteinase inhibitor DE-4 (DE4).
DE
OS
     Lonchocarpus capassa (Apple-leaf).
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicotyledons;
```

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rosids; eurosids I; Fabales; Fabaceae; Papilionoideae; Millettieae;
OC
OC
    Lonchocarpus.
    NCBI TaxID=3926;
OX
RN
    [1]
RP
    PROTEIN SEQUENCE.
RC
    TISSUE=Seed;
RA
    Joubert F.J.;
    "Proteinase inhibitors from Lonchocarpus capassa (apple-leaf) seed.";
RT
    Phytochemistry 23:957-961(1984).
RL
    -!- SIMILARITY: Belongs to the Bowman-Birk serine protease inhibitor
CC
CC
        family.
    _____
CC
CC
    This Swiss-Prot entry is copyright. It is produced through a collaboration
CC
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
    the European Bioinformatics Institute. There are no restrictions on its
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    use as long as its content is in no way modified and this statement is not
CC
CC
    removed.
    _____
CC
DR
    HSSP; P01062; 1DF9.
    InterPro; IPR000877; Prot_inh_BBI.
DR
    Pfam; PF00228; Bowman-Birk leg; 2.
DR
    SMART; SM00269; BowB; 1.
DR
    PROSITE; PS00281; BOWMAN BIRK; 1.
DR
    Direct protein sequencing; Protease inhibitor;
KW
KW
    Serine protease inhibitor.
FT
                      26
                               Reactive bond for trypsin (By
    SITE
                25
FT
                               similarity).
    SITE
                               Reactive bond for chymotrypsin (By
FT
                52
                       53
FT
                               similarity).
                18
                      71
                               By similarity.
FT
    DISULFID
                19
                      33
                               By similarity.
FT
    DISULFID
FT
                22
                      67
                              By similarity.
    DISULFID
                      31
                23
                               By similarity.
FT
    DISULFID
    DISULFID
                41
                     48
                               By similarity.
FT
FT
    DISULFID
                45
                       60
                              By similarity.
                     58
                              By similarity.
FT
    DISULFID
                50
    SEQUENCE 80 AA; 8806 MW; 6E8DF76866B871C9 CRC64;
SO
                       45.5%; Score 46; DB 1; Length 80;
 Ouery Match
 Best Local Similarity 37.5%; Pred. No. 60;
          6; Conservative 4; Mismatches
                                            6; Indels
                                                          0; Gaps
                                                                      0;
 Matches
          2 EAAAREACCRECCARA 17
Qу
             |: : : || || |:
Db
         11 ESESSKPCCSSCCTRS 26
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Search completed: December 8, 2005, 16:07:48 Job time : 233 secs

# GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: December 8, 2005, 15:48:56; Search time 37 Seconds

(without alignments)

44.208 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : PIR\_80:\*

1: pir1:\*
2: pir2:\*
3: pir3:\*

4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

Result		% Query				
No.	Score	Match	Length	DB	ID	Description
1	53	52.5	157	2	S36538	E6 protein - human
2	49	48.5	115	2	A36113	antileukoproteinas
3	47.5	47.0	676	2	G84663	hypothetical prote
4	47	46.5	156	1	W6WL47	E6 protein - human
5	46	45.5	166	2	S36485	E6 protein - human
6	46	45.5	191	2	I46412	keratin KAP5.4 - s
7	46	45.5	465	2	S05311	indoleacetamide hy
8	46	45.5	498	2	A48203	interleukin-14 pre
9	46	45.5	571	2	S69210	protein kinase cak
10	46	45.5	1430	2	T34516	hypothetical prote
11	45	44.6	61	2	E82580	hypothetical prote
12	45	44.6	155	1	W6WL8	E6 protein - human
13	45	44.6	157	1	W6WL5	E6 protein - human

23         44         43.6         299         2         C97102           24         44         43.6         370         1         S57347           25         44         43.6         374         1         S50193           26         44         43.6         496         2         F75257           27         44         43.6         1001         1         PWRBFC           28         44         43.6         1001         1         PWRBFC           29         44         43.6         1001         1         PWRBFC           29         44         43.6         1121         2         S30862           30         43.5         43.1         126         2         I46489           31         43         42.6         169         1         S18946           32         43         42.6         217         2         T33353           33         43         42.6         221         2         C34768           34         42.6         399         2         B24698           37         43         42.6         399         2         D96714           39	14 15 16 17 18 19 20 21	45 45 44 44 44 44	44.6 44.6 43.6 43.6 43.6 43.6 43.6	157 273 369 161 186 188 204 251 254	1 2 2 2 2 2 2 2 2 2	W6WLB5 A43862 G75460 S36491 A45910 JC6547 T08072 AH3413 B84901
25         44         43.6         374         1         S50193           26         44         43.6         496         2         F75257           27         44         43.6         994         2         A48849           28         44         43.6         1001         1         PWRBFC           29         44         43.6         1121         2         S30862           30         43.5         43.1         126         2         146489           31         43         42.6         217         2         T33353           33         43         42.6         221         2         C34768           34         43.6         221         2         C34768           34         43         42.6         233         2         S67947           35         43         42.6         399         2         B24698           36         43         42.6         689         2         T08988           37         43         42.6         711         2         A85352           38         43         42.6         711         2         A85352           40	23	44				
27       44       43.6       994       2       A48849         28       44       43.6       1001       1       PWRBFC         29       44       43.6       1121       2       S30862         30       43.5       43.1       126       2       I46489         31       43       42.6       217       2       T33353         33       43       42.6       221       2       C34768         34       43       42.6       233       2       S67947         35       43       42.6       689       2       T08988         36       43       42.6       689       2       T08988         37       43       42.6       711       2       A85352         38       43       42.6       711       2       A85352         38       43       42.6       121       2       JC6548         41 <t< td=""><td></td><td></td><td></td><td></td><td></td><td>S50193</td></t<>						S50193
28       44       43.6       1001       1       PWRBFC         29       44       43.6       1121       2       S30862         30       43.5       43.1       126       2       I46489         31       43       42.6       169       1       S18946         32       43       42.6       217       2       T33353         33       43       42.6       221       2       C34768         34       43       42.6       399       2       B24698         36       43       42.6       689       2       T08988         37       43       42.6       689       2       T08988         37       43       42.6       689       2       D96714         39       42.5       42.1       931       2       H96527         40       42       41.6       122       2       JC6548         41       42       41.6       223       2       B38346         42       42.1       230       2       A38346         43       42       41.6       230       2       A38346         43       42						
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32       43       42.6       217       2       T33353         33       43       42.6       221       2       C34768         34       43       42.6       233       2       S67947         35       43       42.6       399       2       B24698         36       43       42.6       689       2       T08988         37       43       42.6       976       2       D96714         39       42.5       42.1       931       2       H96527         40       42       41.6       122       2       JC6548         41       42       41.6       122       2       JC6548         41       42       41.6       223       2       B38346         42       42.1       41.6       230       2       A38346         43       42       41.6       237       2       C86452         45       42       41.6       327       2       C86452         45       42       41.6       1212       2       B82809         46       42       41.6       1212       2       B82809         46						
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35       43       42.6       399       2       B24698         36       43       42.6       689       2       T08988         37       43       42.6       711       2       A85352         38       43       42.6       976       2       D96714         39       42.5       42.1       931       2       H96527         40       42       41.6       122       2       JC6548         41       42       41.6       223       2       B38346         42       42       41.6       230       2       A38346         42       42       41.6       230       2       A38346         43       42       41.6       230       2       A38346         43       42       41.6       237       2       C86452         45       42       41.6       327       2       C86452         45       42       41.6       237       2       T16881         47       41       40.6       67       2       T37199         48       41       40.6       164       2       T24272         50       4						
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68 40 39.6 174 2 S71554 69 40 39.6 181 2 A86451	66		39.6	132	1	TIHUSP
69 40 39.6 181 2 A86451						
70 40 39.6 264 2 JC6125	70	40	39.6	264	2	JC6125

E6 protein - human 29K peripheral mem hypothetical prote E6 protein - human ultra-high-sulfur high sulfur protei proteinase inhibit nitrogen fixation hypothetical prote hypothetical prote Ca2+/calmodulin-de Ca2+/calmodulin-de hypothetical prote Ca2+-transporting Ca2+-transporting DNA dependent ATPa cysteine-rich hair ultra high-sulfur hypothetical prote ORF2 protein - Orf alkyl hydroperoxid formate dehydrogen cadmium-transporti cadmium-transporti DNA-directed RNA p protein F27J15.16 high sulfur protei ultra-high-sulfur ultra-high-sulfur hypothetical prote protein F6N18.11 [ exodeoxyribonuclea hypothetical prote hypothetical prote hair keratin cyste hypothetical prote hypothetical prote hypothetical prote hypothetical prote probable endonucle protein F12F1.7 [i cellulose 1,4-beta iron-sulfur cluste hypothetical prote hypothetical prote probable C2H2-type glutamate receptor glucose starvation protein YDR034w-b hypothetical prote allergenic protein hypothetical prote antileukoproteinas hypothetical prote pathogenesis-relat probable ferredoxi U2 small nuclear r

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# ALIGNMENTS

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S36538
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C; Accession: S36538
R; Delius, H.; Hofmann, B.
submitted to the EMBL Data Library, August 1993
A; Description: Primer-directed sequencing of human papillomavirus types.
A; Reference number: S36469
A; Accession: S36538
A; Molecule type: DNA
A; Residues: 1-157 < DEL>
A;Cross-references: UNIPROT:P36803; UNIPARC:UPI00001383B8; EMBL:X74466;
NID:g396910; PIDN:CAA52496.1; PID:g396911
C; Superfamily: papillomavirus E6 protein
C; Keywords: DNA binding; early protein; nucleus; zinc finger
  Ouery Match
                          52.5%; Score 53; DB 2; Length 157;
  Best Local Similarity
                          53.3%; Pred. No. 2.9;
  Matches
            8; Conservative
                                 1; Mismatches
                                                   6; Indels
                                                                  0; Gaps
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1 WEAAAREACCRECCA 15
Qу
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           63 WKGHFVTACCRSCCA 77
Db
RESULT 2
A36113
antileukoproteinase precursor - pig
C; Species: Sus scrofa domestica (domestic pig)
C;Date: 28-Mar-1991 #sequence revision 13-Jan-1993 #text change 09-Jul-2004
C; Accession: A36113; A49198
R; Farmer, S.J.; Fliss, A.E.; Simmen, R.C.M.
Mol. Endocrinol. 4, 1095-1104, 1990
A; Title: Complementary DNA cloning and regulation of expression of the messenger
RNA encoding a pregnancy-associated porcine uterine protein related to human
antileukoproteinase.
A; Reference number: A36113; MUID: 91155942; PMID: 2293019
A; Accession: A36113
A; Status: preliminary
A; Molecule type: mRNA
A; Residues: 1-115 <FAR>
A;Cross-references: UNIPROT:P22298; UNIPARC:UPI0000125858; GB:M57446;
NID:g164319; PIDN:AAA63446.1; PID:g164320
A; Note: the authors translated the codon GCT for residue 52 as Gly
R; Simmen, R.C.; Michel, F.J.; Fliss, A.E.; Smith, L.C.; Fliss, M.F.
Endocrinology 130, 1957-1965, 1992
A; Title: Ontogeny, immunocytochemical localization, and biochemical properties
of the pregnancy-associated uterine elastase/cathepsin-G protease inhibitor,
antileukoproteinase (ALP): monospecific antibodies to a synthetic peptide
recognize native ALP.
A; Reference number: A49198; MUID: 92191891; PMID: 1547723
A; Accession: A49198
A; Status: preliminary
A; Molecule type: protein
A; Residues: 9-26 <SIM>
A; Cross-references: UNIPARC: UPI0000087C99
A; Experimental source: uterus
A; Note: sequence extracted from NCBI backbone (NCBIP:89471)
C; Superfamily: antileukoproteinase; antileukoproteinase repeat homology
F;14-59/Domain: antileukoproteinase repeat homology <ALP1>
F;68-113/Domain: antileukoproteinase repeat homology <ALP2>
                          48.5%; Score 49; DB 2; Length 115;
  Query Match
                          40.0%; Pred. No. 8;
  Best Local Similarity
             6: Conservative
                               4; Mismatches
                                                    5; Indels
                                                                  0; Gaps
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            1 WEAAAREACCRECCA 15
Qу
              |: :: |||: ||
           38 WQCPDKKKCCRDTCA 52
Db
RESULT 4
W6WL47
E6 protein - human papillomavirus type 47
C; Species: human papillomavirus type 47
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A; Note: host Homo sapiens (man)

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C;Date: 31-Mar-1991 #sequence revision 31-Mar-1991 #text change 09-Jul-2004
C; Accession: A35324
R; Kiyono, T.; Adachi, A.; Ishibashi, M.
Virology 177, 401-405, 1990
A; Title: Genome organization and taxonomic position of human papillomavirus type
47 inferred from its DNA sequence.
A; Reference number: A35324; MUID: 90281611; PMID: 2162112
A; Accession: A35324
A; Status: translation not shown
A; Molecule type: DNA
A; Residues: 1-156 <KIY>
A; Cross-references: UNIPROT: P22422; UNIPARC: UPI00001383D9; GB: M32305;
NID:q333062; PIDN:AAA46976.1; PID:g333064
C: Superfamily: papillomavirus E6 protein
C: Keywords: DNA binding; early protein; transforming protein; zinc finger
F;40-76/Region: zinc finger CCCC motif
F;113-149/Region: zinc finger CCCC motif
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  Query Match
                         46.7%; Pred. No. 18;
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Qу
              |: : |||| ||:
Db
           64 WKDYSVYACCRLCCS 78
RESULT 6
I46412
keratin KAP5.4 - sheep
C:Species: Ovis orientalis aries, Ovis ammon aries (domestic sheep)
C; Date: 16-Aug-1996 #sequence revision 16-Aug-1996 #text_change 09-Jul-2004
C; Accession: I46412; S34215
R; Jenkins, B.J.; Powell, B.C.
J. Invest. Dermatol. 103, 310-317, 1994
A; Title: Differential expression of genes encoding a cysteine-rich keratin
family in the hair cuticle.
A; Reference number: I46412; MUID: 94358466; PMID: 7521375
A:Accession: I46412
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 1-191 <JEN>
A; Cross-references: UNIPROT: Q28583; UNIPARC: UPI0000088B92; EMBL: X73434;
NID:q313719; PIDN:CAA51829.1; PID:q313720
C; Genetics:
A; Gene: KRTAP5.4
C; Superfamily: ultra-high-sulfur keratin
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                          38.5%; Pred. No. 28;
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                                                   2; Indels
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           5; Conservative
            5 AREACCRECCARA 17
Qу
              ::::||| ||:::
          167 SOSSCCRPCCSOS 179
Search completed: December 8, 2005, 16:08:31
Job time : 40 secs
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# GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: December 8, 2005, 16:07:58; Search time 12 Seconds

(without alignments)

7.911 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 32527 segs, 5584426 residues

Total number of hits satisfying chosen parameters: 32527

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : Published\_Applications\_AA\_New:\*

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4: /cgn2\_6/ptodata/1/pubpaa/US08\_NEW\_PUB.pep:\*
5: /cgn2\_6/ptodata/1/pubpaa/PCT\_NEW\_PUB.pep:\*

6: /cgn2\_6/ptodata/1/pubpaa/US10\_NEW\_PUB.pep:\*
7: /cgn2\_6/ptodata/1/pubpaa/US11\_NEW\_PUB.pep:\*

8: /cgn2\_6/ptodata/1/pubpaa/US60\_NEW\_PUB.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

7
, Appl
Appli
, Appl
Appli
, Appl

10	40	39.6	321	6	US-10-478-345-8	Sequence 8, Appli
11	39.5	39.1	898	7	US-11-174-150-43	Sequence 43, Appl
12	39	38.6	211	6	US-10-821-234-1372	Sequence 1372, Ap
13	39	38.6	998	6	US-10-510-524-1	Sequence 1, Appli
14	38	37.6	544	6	US-10-980-388-40	Sequence 40, Appl
15	38	37.6	831	6	US-10-467-657-4486	Sequence 4486, Ap
16	38	37.6	1907	7	US-11-039-398-25	Sequence 25, Appl
17	37.5	37.1	247	6	US-10-632-150 <b>-</b> 36	Sequence 36, Appl
18	37.5	37.1	247	7	US-11-073-457-36	Sequence 36, Appl
19	37.5	37.1	511	7	US-11-012-762-66	Sequence 66, Appl
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21	37	36.6	145	6	US-10-467-657-4246	Sequence 4246, Ap
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24	37	36.6	376	7	US-11-116-939-8	Sequence 8, Appli
25	37	36.6	575	6	US-10-980-388-46	Sequence 46, Appl
26	36.5	36.1	1036	6	US-10-131-826A-142	Sequence 142, App
27	36	35.6	336	6	US-10-478-345-4	Sequence 4, Appli
28	36	35.6	393	6	US-10-821-234-1292	Sequence 1292, Ap
29	36	35.6	548	7	US-11-137-465-47	Sequence 47, Appl
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35	35	34.7	263	6	US-10-131-826A-484	Sequence 484, App
36	35	34.7	263	6	US-10-821-234-1403	Sequence 1403, Ap
37	35	34.7	346	6	US-10-878-556A-121	Sequence 121, App
38	35	34.7	346	7	US-11-069-642-109	Sequence 109, App
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47	34	33.7	337	6	US-10-467-962B-97	Sequence 97, Appl
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49	34	33.7	816	7	US-11-090-439-48	Sequence 48, Appl
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56	33.5	33.2	1028	7	US-11-067-121-7	Sequence 7, Appli
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63	33	32.7	148	6	US-10-526-716-2	Sequence 2, Appli
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71	33	32.7	776	6	US-10-925-970-3	Sequence	3, Appli
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88	32	31.7	101	7	US-11-082-381-1	Sequence	1, Appli
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# ALIGNMENTS

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; Publication No. US20050244435A1
; GENERAL INFORMATION:
; APPLICANT: Shone, Clifford Charles
; APPLICANT: Quinn, Conrad Padraig
; APPLICANT: Foster, Keith Alan
; APPLICANT: Chaddock, John
; APPLICANT: Marks, Philip
; APPLICANT: Sutton, J. Mark
; APPLICANT: Stancombe, Patrick
; APPLICANT: Wayne, Jonathan
; TITLE OF INVENTION: Recombinant Toxin Fragments
; FILE REFERENCE: 1581.0130004
; CURRENT APPLICATION NUMBER: US/11/077,550
; CURRENT FILING DATE: 2005-03-11
; PRIOR APPLICATION NUMBER: 10/241,596
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RESULT 1

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  PRIOR APPLICATION NUMBER: 09/255,829
  PRIOR FILING DATE: 1999-02-23
  PRIOR APPLICATION NUMBER: PCT/GB97/02273
  PRIOR FILING DATE: 1997-08-22
  PRIOR APPLICATION NUMBER: 08/782,893
  PRIOR FILING DATE: 1996-12-27
  PRIOR APPLICATION NUMBER: GB9625996.5
  PRIOR FILING DATE: 1996-12-13
  PRIOR APPLICATION NUMBER: GB9617671.4
; PRIOR FILING DATE: 1996-08-23
; NUMBER OF SEQ ID NOS: 179
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; Sequence 2, Application US/11085775
; Publication No. US20050260634A1
; GENERAL INFORMATION:
; APPLICANT: BALDWIN, DARYL
; APPLICANT: CLARK, HILARY
  APPLICANT: JUBB, ADRIAN
; APPLICANT: KOEPPEN, HARTMUT
; APPLICANT: QUAN, CLIFFORD
; APPLICANT: WU, THOMAS
; APPLICANT: ZHANG, ZEMIN
  TITLE OF INVENTION: ACHAETE-SCUTE LIKE-2 POLYPEPTIDES AND ENCODING NUCLEIC
  TITLE OF INVENTION: ACIDS AND METHODS FOR THE DIAGNOSIS AND TREATMENT OF
TUMOR
  FILE REFERENCE: P5028R1P1-US
  CURRENT APPLICATION NUMBER: US/11/085,775
  CURRENT FILING DATE: 2005-03-21
  PRIOR APPLICATION NUMBER: PCT/US03/17682
  PRIOR FILING DATE: 2003-06-04
  PRIOR APPLICATION NUMBER: US 10/454,945
  PRIOR FILING DATE: 2003-06-04
  PRIOR APPLICATION NUMBER: US 60/407,087
  PRIOR FILING DATE: 2002-08-29
; NUMBER OF SEQ ID NOS: 78
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   LENGTH: 3500
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US-11-085-775-2

Db

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Search completed: December 8, 2005, 16:19:27 Job time : 13 secs

1274 AAAAACCAGCC 1284

# GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: December 8, 2005, 16:08:38; Search time 162 Seconds

(without alignments)

43.846 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 1867569 segs, 417829326 residues

Total number of hits satisfying chosen parameters: 1867569

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

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	4	101	100.0	17	3	US-09-813-197-4	Sequence 4, Appli
	5	101	100.0	17	4	US-10-126-752 <b>-1</b>	Sequence 1, Appli
	6	101	100.0	17	4	US-10-174-368A-3	Sequence 3, Appli
	7	101	100.0	17	4	US-10-345-281-48	Sequence 48, Appl
	8	101	100.0	17	4	US-10-264-127-4	Sequence 4, Appli
	9	101	100.0	17	4	US-10-339-712-4	Sequence 4, Appli
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; Sequence 3, Application US/09973145
; Patent No. US20020132248A1
; GENERAL INFORMATION:
  APPLICANT: Rothschild, Kenneth J.
  APPLICANT:
              Gite, Sadanand
  APPLICANT:
              Olejnik, Jerzy
  TITLE OF INVENTION: N-Terminal and C-Terminal Markers in Nascent Proteins
   FILE REFERENCE: AMBER-06819
   CURRENT APPLICATION NUMBER: US/09/973,145
   CURRENT FILING DATE:
                         2001-10-09
   PRIOR APPLICATION NUMBER: 09/382,950
   PRIOR FILING DATE: 1999-08-25
   NUMBER OF SEQ ID NOS: 18
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; SEQ ID NO 3
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Search completed: December 8, 2005, 16:22:17

Job time : 164 secs

# GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: December 8, 2005, 15:49:22; Search time 46 Seconds

(without alignments)

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Title: US-10-772-164-1

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Listing first 100 summaries

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    APPLICANT: Tsien, Roger Y.
    APPLICANT: Griffin, B. Albert
    TITLE OF INVENTION: TARGET SEQUENCES FOR SYNTHETIC MOLECULES
    NUMBER OF SEQUENCES: 4
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Fish & Richardson P.C.
      STREET: 4225 Executive Square, Suite 1400
      CITY: La Jolla
      STATE: CA
      COUNTRY: USA
      ZIP: 92037
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: Windows 95
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SOFTWARE: FastSEQ for Windows Version 2.0b
     CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/955,206
       FILING DATE: 21-OCT-1997
    ATTORNEY/AGENT INFORMATION:
    NAME: Haile, Ph.D., Lisa A.
      REGISTRATION NUMBER: 38,347
     REFERENCE/DOCKET NUMBER: 07257/060001
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 619/678-5070
       TELEFAX: 619/678-5099
  INFORMATION FOR SEQ ID NO: 1:
   SEQUENCE CHARACTERISTICS:
      LENGTH: 17 amino acids
       TYPE: amino acid
       TOPOLOGY: linear
    MOLECULE TYPE: peptide
     FEATURE:
      OTHER INFORMATION: the N-terminus is acetylated and
      OTHER INFORMATION: the C-terminus is amidated
US-08-955-206-1
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Qу
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Search completed: December 8, 2005, 16:09:21

Job time : 48 secs

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OM protein - protein search, using sw model

Run on: December 8, 2005, 15:04:13; Search time 184 Seconds

(without alignments)

40.595 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2443163 segs, 439378781 residues

Total number of hits satisfying chosen parameters: 2443163

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : A\_Geneseq\_21:\*

1: geneseqp1980s:\*

2: geneseqp1990s:\*

3: geneseqp2000s:\*

4: geneseqp2001s:\*

5: geneseqp2002s:\*

6: geneseqp2003as:\*

7: geneseqp2003bs:\*

8: geneseqp2004s:\*

o. genebedp2001b.

9: geneseqp2005s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

### SUMMARIES

,	Result		% Query						
,	No.	Score	Match Length DB			ID	Descripti	Description	
	1	101	100.0	17	2	AAY05336	Aay05336	Target se	
	2	101	100.0	17	3	AAB20847	Aab20847	Peptide a	
	3	101	100.0	17	4	AAB35430	Aab35430	Dye-bindi	
	4	101	100.0	17	4	AAM48100	Aam48100	Fluoresce	
	5	101	100.0	17	8	AD006947	Ado06947	FLASH-bin	
	6	101	100.0	17	9	ADZ76895	Adz76895	RNA-tag f	
	7	90	89.1	17	2	AAY05337	Aay05337	Target se	
	8	90	89.1	17	3	AAB20848	Aab20848	Peptide a	

9	87	86.1	19	4	AAM51838	Aam51838	Gene corr
10	87	86.1	19	5	AAU81286		Plasmid e
11	87	86.1	19	5	AAU75749		FLAsH pep
12	87	86.1	19	7	ADB78479		FIAsH pep
13	81	80.2	19	7	ABR84531		. FLAsH pep
14	76	75.2	595	8	ADQ76865		Adenosine
15	61	60.4	22	3	AAY88739		Core poly
16	61	60.4	22	4	AAB77094		Core poly
17	61	60.4	22	4	ABB00098		Viral DP1
18	61	60.4	22	4	AAU12647		DP178-lik
19	61	60.4	55	5	ADE01583		Hybrid po
20	56.5	55.9	106	7	ABO76100		Pseudomon
21	53	52.5	365	7	ABO83225		Pseudomon
22	52	51.5	631	7	AB071317		Pseudomon
23	51	50.5	535	8	ADL70535		Human G-p
24	50	49.5	28	3	AAY88872		Core poly
25	50	49.5	28	4	AAB77227		Core poly
26	50	49.5	28	4	ABB00231		. Viral DP1
27	50	49.5	28	4	ABB01704		Viral cor
28	50	49.5	28	4	AAU12780		DP178-lik
29	50	49.5	28	6	ABO10317		HIV-1 BRU
30	50	49.5	30	8	ADT71522		Linker mo
31	50	49.5	32	8	ADT71523		Linker mo
32	50	49.5	35	8	ADT71524		Linker mo
33	50	49.5	62	7	ABO80197		Pseudomon
34	50	49.5	906	8	ADP31344		Human sec
35	50	49.5	1134	8	ADP30647		Human sec
36	49.5	49.0	161	7	ABO79455	_	Pseudomon
37	49	48.5	113	7	ABO71027	Abo71027	Pseudomon
38	49	48.5	120	2	AAW07542		Clone 99,
39	49	48.5	918	8	ADP31459		Human sec
40	49	48.5	1626	8	ADP31008		Human sec
41	48	47.5	126	2	AAW98909		Mouse IMC
42	48	47.5	131	2	AAW98908	Aaw98908	Mouse IMC
43	48	47.5	131	7	ADE25527	Ade25527	Mouse SLP
44	48	47.5	131	7	ADF28912	Adf28912	Mouse SLP
45	48	47.5	131	9	ADX02863	Adx02863	Murine an
46	48	47.5	146	8	ADQ59487	Adq59487	Human can
47	48	47.5	146	9	ADZ13856	Adz13856	Murine ca
48	48	47.5	162	7	ABO81835	Abo81835	Pseudomon
49	48	47.5	1305	8	ADP31389	Adp31389	Human sec
50	48	47.5	1312	8	ADP30999	Adp30999	Human sec
51	48	47.5	2001	8	ADP31644	Adp31644	Human sec
52	48	47.5	2260	8	ADP30687	Adp30687	Human sec
53	48	47.5	2272	8	ADP31136	Adp31136	Human sec
54	48	47.5	4440	6	ABU88256	Abu88256	Novel hum
55	48	47.5	4440	6	ABU90135	Abu90135	Novel hum
56	48	47.5	4440	6	ABU96437	Abu96437	Novel hum
57	48	47.5	4440	6	ABU99046	Abu99046	Novel hum
58	48	47.5	4440	6	ABU98261	Abu98261	. Novel hum
59	48	47.5	4440	6	ABU91967		Novel hum
60	48	47.5	4440	6	ABU85271		Novel hum
61	48	47.5	4440	6	ABO00410		Novel hum
62	48	47.5	4440	6	ABU88961		Novel hum
63	48	47.5	4440	6	ABO06457		Novel hum
64	48	47.5	4440	6	ABU95517		Novel hum
65	48	47.5	4440	6	ABU95207	Abu95207	Novel hum

```
Abu90755 Novel hum
                 4440 6 ABU90755
66
       48
            47.5
            47.5 4440 6 ABU93917
                                                   Abu93917 Novel hum
67
       48
                                                   Abu86191 Novel hum
       48
            47.5 4440 6 ABU86191
68
                                                   Abu82046 Novel hum
            47.5
                  4440 6
                           ABU82046
69
       48
                                                   Abu07907 Novel hum
70
       48
            47.5
                  4440 6
                           ABU07907
                 4440 6
                                                   Abu94227 Novel hum
71
       48
            47.5
                           ABU94227
                                                   Abo00100 Novel hum
            47.5 4440 6 ABO00100
72
       48
                                                  Abu87111 Novel hum
            47.5 4440 6 ABU87111
73
       48
            47.5 4440 6 ABU91352
                                                  Abu91352 Novel hum
74
        48
            47.5 4440 6 ABU90445
                                                  Abu90445 Novel hum
75
       48
                                                  Abu97036 Novel hum
                 4440 6 ABU97036
76
            47.5
        48
                 4440 6 ABO05232
                                                   Abo05232 Novel hum
77
            47.5
       48
                                                   Ade25528 Rat SLPI
78
       47
            46.5
                   131 7 ADE25528
                   131 7 ADF28911
                                                   Adf28911 Rat SLPI
79
       47
            46.5
                   170 8 ADS10852
                                                   Ads10852 Human the
80
       47
            46.5
       47
            46.5
                   195 8 ADP30696
                                                   Adp30696 Human sec
81
                   205 8 ADS10854
       47
            46.5
                                                   Ads10854 Human the
82
            46.5
                    210 9 AEA15447
                                                   Aea15447 Human pol
83
        47
                    357 8 ADP31267
                                                   Adp31267 Human sec
84
       47
            46.5
                                                   Adp31147 Human sec
85
        47 46.5
                   621 8 ADP31147
86
        47 46.5
                   783 8 ADP31436
                                                   Adp31436 Human sec
87
        47
            46.5
                 821 8 ADP30679
                                                   Adp30679 Human sec
        47
            46.5
                   821 8 ADP30680
                                                   Adp30680 Human sec
88
                   882 8 ADP31688
                                                   Adp31688 Human sec
89
       47
            46.5
90
            46.5
                   990 8 ADP31553
                                                   Adp31553 Human sec
       47
91
       47
            46.5 1033 8 ADP30984
                                                   Adp30984 Human sec
       47
            46.5 1224 8 ADP31426
                                                   Adp31426 Human sec
92
       47 46.5 1518 8 ADP31532
                                                   Adp31532 Human sec
93
            46.5 1665 8 ADP31187
                                                   Adp31187 Human sec
94
       47
            46.5 1679 4 AAU07343
                                                   Aau07343 1-aminocy
95
        47
                   2058 8 ADP31630
                                                   Adp31630 Human sec
96
        47
            46.5
97
        47
            46.5
                   2187 8 ADP30882
                                                   Adp30882 Human sec
98
        47
            46.5
                   3201 8 ADP31545
                                                   Adp31545 Human sec
                                                   Adp31148 Human sec
99
        47
            46.5
                   3390 8 ADP31148
100
        47
            46.5
                   3411 8 ADP30667
                                                   Adp30667 Human sec
```

#### ALIGNMENTS

```
AAY05336
     AAY05336 standard; peptide; 17 AA.
ID
XX
AC
     AAY05336;
XX
DΤ
     29-JUN-1999 (first entry)
XX
DE
     Target sequence peptide, SEQ ID NO. 1.
XX
     Biarsenical compound; alpha-helix peptide; polypeptide purification;
KW
KW
     immunoassay; crosslinking agent.
XX
OS
     Synthetic.
XX
PN
     W09921013-A1.
XX
```

RESULT 1

```
PD
     29-APR-1999.
XX
PF
                    98WO-US022363.
     21-OCT-1998;
XX
PR
     21-OCT-1997;
                    97US-00955050.
PR
     21-OCT-1997;
                   97US-00955206.
                   97US-00955859.
PR
     21-OCT-1997;
XX
     (REGC ) UNIV CALIFORNIA.
PΑ
XX
     Tsien RY, Griffin AB;
PΙ
XX
DR
    WPI; 1999-288410/24.
XX
PT
     Biarsenical compounds that react specifically with cysteine residues.
XX
PS
     Claim 10; Page 41; 77pp; English.
XX
CC
     This sequence represents a target alpha-helix sequence for the
     biarsenical compounds (BC) of the invention, which are able to react
CC
CC
     specifically with cysteine residues in a target sequence to generate a
CC
     detectable signal. The BCs are used: (i) as labels that allow
CC
     identification of carrier molecules, e.g. in polypeptide purification,
CC
     immunoassays or other chemical or biological assays, including labelling
CC
     in vivo, e.g. to identify, locate or quantify polypeptides or nucleic
CC
     acids); (ii) for attaching a polypeptide to a solid substrate; or (iii)
     to induce a polypeptide domain to adopt a more nearly alpha-helical form,
CC
CC
     e.g. a conformation that can bind a drug. Tetra-arsenical compounds
     derived from the BCs are used to crosslink two binding partners, e.g. to
CC
CC
     study the effect of dimerisation on signal transduction. The BCs react
CC
     specifically with Cys-containing targets, and can be engineered to have
CC
     particular properties, especially ability to cross a biological membrane
CC
     and absence of any self-fluorescence. Both the BC and its target sequence
     are small, and BC binding between them is reversible, e.g. by treatment
CC
CC
     with a dithiol. Particularly, the BC becomes fluorescent when bound to
CC
     its target, but with a significant red-shift from the fluorescence of
CC
     fluorescein, allowing detection with very low background
XX
SO
     Sequence 17 AA;
  Query Match
                          100.0%; Score 101; DB 2;
                                                     Length 17;
  Best Local Similarity
                          100.0%; Pred. No. 1.7e-05;
  Matches
           17; Conservative
                               0; Mismatches
                                                   0; Indels
                                                                 0; Gaps
                                                                             0;
Qу
            1 WEAAAREACCRECCARA 17
              Db
           1 WEAAAREACCRECCARA 17
RESULT 2
AAB20847
     AAB20847 standard; peptide; 17 AA.
XX
AC
    AAB20847;
XX
DT
     03-JAN-2001 (first entry)
XX
```

```
DE
     Peptide amino acid sequence SEQ ID NO:48.
XX
KW
    Target protein binding element; protein level control; eukaryotic;
    ubiquitination recognition element; treatment; infection; cancer;
KW
     inflammatory condition; genetic disease; insecticide; herbicide;
KW
    antiviral; antiparasitic; hepatotropic; antiinflammatory; cytostatic;
KW
    tumour; pest control; pesticide; rodenticide; fungicide; gene expression;
KW
KW
    gene therapy.
XX
OS
    Unidentified.
XX
    WO200047220-A1.
PN
XX
PD
    17-AUG-2000.
XX
PF
    11-FEB-2000; 2000WO-US003436.
XX
PR
    12-FEB-1999;
                   99US-0119851P.
PR
    28-SEP-1999;
                   99US-00406781.
XX
     (PROT-) PROTEINIX INC.
PΑ
XX
ΡI
    Kenten JH, Roberts SF, Lebowitz MS;
XX
DR
    WPI; 2000-565258/52.
XX
    Novel compounds for modulating the ubiquitination of target proteins
PT
     comprising a ubiquitination recognition element-target protein element
PT
     fusion, useful for treating viral infections.
PT
XX
    Disclosure; Page 55; 106pp; English.
PS
XX
     The present invention describes a compound (I) for activating the
CC
    ubiquitination (Ub'n) of a target protein comprising a Ub'n recognition
CC
CC
     (peptide) element (URE) covalently linked to a target protein (peptide)
     element (TPE). (I) can have antiviral, antiparasitic, hepatotropic,
CC
     antiinflammatory and cytostatic activities. The compound of (I) may be
CC
    used to treat a viral infection (especially hepatitis A, B, C or G, HIV-1
CC
    or 2, Herpes, CMV, rabies or Rouse sarcoma virus (RSV)), parasitic
CC
     infection, an infection caused by an eukaryotic organism in a mammal, to
CC
     treat a tumour or to control pests. The compound may also be used to
CC
CC
     screen for target protein binding elements, to develop pesticides (e.g.
     insecticides, rodenticides, fungicides and herbicides) and to control
CC
     gene expression (gene therapy). The present sequence represents an
CC
     example of a peptide which is given in the exemplification of the present
CC
CC
     invention
XX
SO
    Sequence 17 AA;
                          100.0%; Score 101; DB 3; Length 17;
 Query Match
                          100.0%; Pred. No. 1.7e-05;
 Best Local Similarity
 Matches
          17; Conservative 0; Mismatches
                                                 0; Indels
                                                                 0; Gaps
                                                                             0;
           1 WEAAAREACCRECCARA 17
Qу
              Db
           1 WEAAAREACCRECCARA 17
```

```
RESULT 29
ABO10317
    ABO10317 standard; peptide; 28 AA.
ID
XX
AC
    ABO10317;
XX
DT
    19-AUG-2003 (first entry)
XX
DE
    HIV-1 BRU qp41 DP178 based-peptide T234.
XX
     HIV; DP107; DP178; glycoprotein 41; antiviral; virucide; EBV;
KW
     Epstein-Barr virus infection; heptad repeat motif.
KW
XX
     Human immunodeficiency virus; isolate BRU.
OS
os
     Synthetic.
XX
PN
     US6518013-B1.
XX
     11-FEB-2003.
PD
XX
                    95US-00485546.
PF
     07-JUN-1995;
XX
                    93US-00073028.
PR
     07-JUN-1993:
                    94US-00255208.
PR
     07-JUN-1994;
                    94US-00360107.
PR
     20-DEC-1994;
XX
PΑ
     (TRIM-) TRIMERIS INC.
XX
PΙ
     Barney SO, Lambert DM,
                              Petteway SR;
XX
     WPI; 2003-465599/44.
DR
XX
     Inhibiting transmission of Epstein-Barr virus to a cell, by contacting
PT
     the cell with a peptide consisting of a region of Epstein-Barr virus
PT
PT
     protein.
XX
     Example; Fig 49G; 716pp; English.
PS
XX
     The invention relates to inhibiting (M) transmission of an Epstein-Barr
CC
     virus to a cell, comprising contacting the cell with an effective
CC
     concentration of a peptide consisting of a region of 16-39 consecutive
CC
     amino acids of an Epstein-Barr virus protein for an effective period of
CC
     time, where the region is recognised by one or more of ALLMOTI5,
CC
     107x178x4 or PLZIP sequence search motifs, the peptide further comprises
CC
     an amino terminal X, and a carboxy terminal Z in which X comprises an
CC
     amino group, acetyl group, 9-fluorenylmethoxy-carbonyl group, hydrophobic
CC
     group or macromolecular carrier group, and Z comprises a carboxyl group,
CC
     amido group, hydrophobic group, or macromolecular carrier group, and
CC
     fusion of the virus to the cell is inhibited. The peptides were
CC
     identified by analysing the structure/motifs present in the HIV-1
CC
     glycoprotein 41 anti-HIV peptides DP107 and DP178. These heptad repeat
CC
     motif containing peptides were used to design the motifs cited above,
CC
     which in turn were used to analyse proteins from other pathogenic
CC
     organisms and HIV isolates, looking for DP107/178 structural analogues.
CC
     The method is useful for inhibiting transmission of Epstein-Barr virus to
CC
```

```
a cell and Epstein-Barr virus infection. The present sequence is a
CC
    antiviral peptide based on a region of a protein from a pathogenic
CC
CC
    organism analogous to DP107 or DP178
XX
    Sequence 28 AA;
SO
                         49.5%; Score 50; DB 6; Length 28;
 Query Match
 Best Local Similarity 73.3%; Pred. No. 33;
          11; Conservative 0; Mismatches
                                                4; Indels
 Matches
                                                                 0; Gaps
                                                                             0;
           2 EAAAREACCRECCAR 16
Qу
              1 EAAAREAAAREAAAR 15
Db
RESULT 38
AAW07542
    AAW07542 standard; protein; 120 AA.
ID
XX
AC
    AAW07542;
XX
DT
    07-FEB-1997 (first entry)
XX
DĖ
    Clone 99, human pro-opiomelanocortin cDNA analogue protein prod. (2).
XX
    Human; poly(A) RNA; cDNA synthesis; polymerase chain reaction;
KW
KW
     lambda gt11; phage vector; PCR; amplification; clone 99;
    pro-opiomelanocortin.
KW
XX
OS
    Homo sapiens.
XX
FH
                     Location/Qualifiers
FT
    Misc-difference 101
FT
                     /note= "corresponding codon TGA"
XX
    EP716150-A1.
PN
XX
PD
    12-JUN-1996.
XX
PF
    05-DEC-1995;
                   95EP-00119121.
XX
PR
     05-DEC-1994;
                   94JP-00300657.
XX
     (TAKE ) TAKEDA CHEM IND LTD.
PΑ
XX
PΙ
    Onda H, Hosoya M;
XX
DR
    WPI; 1996-269991/28.
DR
    N-PSDB; AAT43979.
XX
PT
     DNA primers for sequences encoding Gly-Lys-Arg, Gly-Arg-Arg or Gly-Lys-
PT
     Lys - useful for identifying peptide(s) with useful physiological
PΤ
     activity having the specified sequences at their C-terminal ends.
XX
PS
     Example 3; Fig 10; 37pp; English.
XX
    Human poly(A) RNA was used as a template for cDNA synthesis, conducted by
CC
CC
    using, as primers, antisense codons for Gly-Lys-Arg, Gly-Arg-Arg or Gly-
```

```
Lys-Lys. The prod. was ligated into a lambda gtll phage vector, and PCR
CC
    amplified. The prod. was subcloned with a TA receptor, and cDNA fragments
CC
    from 100 clones sequenced, including clone 99, which was decoded in 3
CC
    reading frames to give AAW07541-43. The nucleotide sequence of clone 99
CC
    was found to have a portion of cDNA encoding human pro-opiomelanocortin,
CC
    an entire sequence of cDNA encoding gamma-MSH and a sequence identical
CC
    with the 5'-upstream region of Gly-Arg-Arg
CC
XX
SO
    Sequence 120 AA;
                        48.5%; Score 49; DB 2; Length 120;
 Query Match
                        64.3%; Pred. No. 1.4e+02;
 Best Local Similarity
          9; Conservative 0; Mismatches 5; Indels
 Matches
                                                              0; Gaps
                                                                          0;
           3 AAAREACCRECCAR 16
Qу
             37 AAARGPCCWPCCFR 50
Db
```

Search completed: December 8, 2005, 16:03:54

Job time : 189 secs